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UNITED STATES DISTRICT COURT
Northern District of California
450 Golden Gate Avenue
San Francisco, California 94102

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Richard W. Wieking
Clerk

General Court Number
415.522.2000

May 23, 2008

Northern District of Illinois
219 South Dearborn Street
Chicago, IL 60604

FILED

JUN 02 2008

RE: CV 08-01070 MEJ LOUIS FUENTES-v-BAYER CORPORATION

MICHAEL W. DOBBINS
CLERK, U.S. DISTRICT COURT

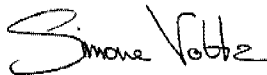
Dear Clerk,

Pursuant to an order transferring the above captioned case to your court, transmitted herewith are:

- ☒ All docket entries/documents.
- ☒ Transferral Order.

Please acknowledge receipt of the above documents on the attached copy of this letter.

Sincerely,
RICHARD W. WIEKING, Clerk



by: Simone Voltz
Case Systems Administrator

Enclosures
Copies to counsel of record

ADRMOP, CLOSED, E-Filing, TRANSF

**U.S. District Court
California Northern District (San Francisco)
CIVIL DOCKET FOR CASE #: 3:08-cv-01070-MEJ
Internal Use Only**

Fuentes v. Bayer Corporation et al
Assigned to: Magistrate Judge Maria-Elena James
Cause: 28:1332 Diversity-Personal Injury

Date Filed: 02/22/2008
Date Terminated: 05/14/2008
Jury Demand: Plaintiff
Nature of Suit: 365 Personal Inj. Prod.
Liability
Jurisdiction: Diversity

Plaintiff

Louis Fuentes

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V.

Defendant

Bayer Corporation

Defendant

Baxter Healthcare Corporation

Defendant

**Armour Pharmaceutical Company
Inc**

Defendant

Alpha Therapeutic Corporation

Date Filed	#	Docket Text
02/22/2008	<u>1</u>	COMPLAINT /Issued summons against Bayer Corporation, Baxter Healthcare Corporation, Armour Pharmaceutical Company Inc, Alpha Therapeutic Corporation (Filing fee \$ 350, receipt number 34611016128.). Filed byLouis Fuentes. (ga, COURT STAFF) (Filed on 2/22/2008) (Additional attachment(s) added on 3/31/2008: # <u>1</u> Civil Cover Sheet) (ga, COURT STAFF). (Entered: 02/26/2008)
02/22/2008	<u>2</u>	ADR SCHEDULING ORDER: Case Management Conference set for

		5/29/2008 10:00 AM. Case Management Statement due by 5/22/2008.. Signed by Judge Maria-Elena James on 2/22/08. (Attachments: # <u>1</u> Judge Standing Order, # <u>2</u> Court Standing Order, # <u>3</u> Consent/Decline Form)(ga, COURT STAFF) (Filed on 2/22/2008) (Entered: 02/26/2008)
02/22/2008		CASE DESIGNATED for Electronic Filing. (ga, COURT STAFF) (Filed on 2/22/2008) (Entered: 02/26/2008)
05/08/2008	<u>3</u>	CERTIFICATE OF SERVICE by Louis Fuentes re <u>1</u> Complaint, <i>Proof of Service Summons & Complaint - Baxter Healthcare Corporation</i> (Foster, Heather) (Filed on 5/8/2008) (Entered: 05/08/2008)
05/08/2008	<u>4</u>	SUMMONS Returned Executed by Louis Fuentes. Bayer Corporation served on 3/14/2008, answer due 4/3/2008. (<i>Proof of Service Summons & Complaint</i>) (Foster, Heather) (Filed on 5/8/2008) (Entered: 05/08/2008)
05/08/2008	<u>5</u>	SUMMONS Returned Executed by Louis Fuentes. Alpha Therapeutic Corporation served on 3/14/2008, answer due 4/3/2008. (<i>Proof of Service Summons & Complaint</i>) (Foster, Heather) (Filed on 5/8/2008) (Entered: 05/08/2008)
05/08/2008	<u>6</u>	SUMMONS Returned Executed by Louis Fuentes. Armour Pharmaceutical Company Inc served on 3/14/2008, answer due 4/3/2008. (<i>Proof of Service Summons & Complaint</i>) (Foster, Heather) (Filed on 5/8/2008) (Entered: 05/08/2008)
05/08/2008	<u>7</u>	SUMMONS Returned Executed by Louis Fuentes. Baxter Healthcare Corporation served on 3/14/2008, answer due 4/3/2008. (<i>Proof of Service Summons & Complaint</i>) (Foster, Heather) (Filed on 5/8/2008) (Entered: 05/08/2008)
05/14/2008	<u>8</u>	ORDER TRANSFERRING CASE to the Northern District of Illinois (per CTO-103). (sv, COURT STAFF) (Filed on 5/14/2008) (Entered: 05/23/2008)
05/23/2008	<u>9</u>	CLERK'S NOTICE of case transferred electronically to the Northern District of Illinois <u>8</u> (sv, COURT STAFF) (Filed on 5/23/2008) (Entered: 05/23/2008)

Inasmuch as no objection is
pending at this time, the
stay is lifted.

MAY - 7 2008

CLERK'S OFFICE
JUDICIAL PANEL ON
MULTIDISTRICT LITIGATION

UNITED STATES JUDICIAL PANEL
on
MULTIDISTRICT LITIGATION

JUDICIAL PANEL ON
MULTIDISTRICT LITIGATION

APR 21 2008

FILED
CLERK'S OFFICE

FILED: MAY 7, 2008

08CV2702 CEM

JUDGE GRADY

MDL No. 986

IN RE: "FACTOR VIII OR IX CONCENTRATE BLOOD
PRODUCTS" PRODUCTS LIABILITY LITIGATION

Louis Fuentes v. Bayer Corp., et al.,)
N.D. California, C.A. No. 3:08-1070 - 08cv2702)

CONDITIONAL TRANSFER ORDER (CTO-103)

On December 7, 1993, the Panel transferred 27 civil actions to the United States District Court for the Northern District of Illinois for coordinated or consolidated pretrial proceedings pursuant to 28 U.S.C. § 1407. See 853 F.Supp. 454 (J.P.M.L. 1993). Since that time, 263 additional actions have been transferred to the Northern District of Illinois. With the consent of that court, all such actions have been assigned to the Honorable John F. Grady.

It appears that the action on this conditional transfer order involves questions of fact that are common to the actions previously transferred to the Northern District of Illinois and assigned to Judge Grady.

Pursuant to Rule 7.4 of the Rules of Procedure of the Judicial Panel on Multidistrict Litigation, 199 F.R.D. 425, 435-36 (2001), this action is transferred under 28 U.S.C. § 1407 to the Northern District of Illinois for the reasons stated in the order of December 7, 1993, and, with the consent of that court, assigned to the Honorable John F. Grady.

This order does not become effective until it is filed in the Office of the Clerk of the United States District Court for the Northern District of Illinois. The transmittal of this order to said Clerk shall be stayed 15 days from the entry thereof. If any party files a notice of opposition with the Clerk of the Panel within this 15-day period, the stay will be continued until further order of the Panel.

FOR THE PANEL:

A CERTIFIED TRUE COPY

MAY - 7 2008

ATTEST: *[Signature]*
FOR THE JUDICIAL PANEL ON
MULTIDISTRICT LITIGATION

A TRUE COPY-ATTESTED
MICHAEL W. DOBBINS, CLERK

By *[Signature]* WILLIE A. HAYNES
DEPUTY CLERK
U.S. DISTRICT COURT, NORTHERN
DISTRICT OF ILLINOIS
DATE: MAY 13, 2008

COPY

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FILED

08 FEB 22 PM 3:35
EDWARD V. BREWER
CLERK, U.S. DISTRICT COURT
NORTHERN DISTRICT OF CALIFORNIA

E-filing

MEJ

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Attorneys for Plaintiffs

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UNITED STATES DISTRICT COURT
NORTHERN DISTRICT OF CALIFORNIA

LOUIS FUENTES, a resident of Junction, Texas

Plaintiff,

v.

BAYER CORPORATION, an Indiana corporation,
successor to CUTTER BIOLOGICAL, a California
Corporation; BAXTER HEALTHCARE
CORPORATION, a Delaware corporation, and its
HYLAND DIVISION; ARMOUR PHARMACEUTICAL
COMPANY, INC., a Delaware corporation and ALPHA
THERAPEUTIC CORPORATION, a California
corporation,

Defendants.

Case No.

**COMPLAINT FOR
DAMAGES AND
INJUNCTIVE RELIEF**

1070

Jury Trial Demanded

- (1) Negligence
- (2) Negligence Per Se
- (3) Fraudulent Omission and Concealment
- (4) Breach of Implied Warranty

I. INTRODUCTION

1. Defendants manufactured blood products known as "Factor VIII" and "Factor IX" for the treatment of hemophilia, and sold these products to people with hemophilia in the United States and worldwide, despite knowledge that the products were manufactured from sick, high-risk donors and/or known to be contaminated with the virus that causes Non-A, Non-B Hepatitis (now known as "Hepatitis C" or "HCV"). Defendants knowingly declined to timely pursue or adopt treatment and manufacturing practices that would have prevented the infection of Plaintiff with HCV, as described in more detail below. Defendants also continued selling old stocks of products they knew to be contaminated with HCV even after they or others had introduced safer products. Plaintiff is a person with hemophilia who contracted HCV through use

1 of Defendants' contaminated products. This complaint describes the factual predicate for
2 Plaintiff's infection: a pattern of foot-dragging, denial, and obfuscation by the pharmaceutical
3 companies on whom his health and well-being depended.

4 2. Defendants manufactured HCV-contaminated blood factor products using
5 human plasma taken from thousands of paid donors, including populations then known to be at
6 high risk of carrying blood-borne diseases, such as urban homosexuals, prisoners, and intravenous
7 drug users. Defendants intentionally recruited urban homosexuals who had a history of viral
8 hepatitis as plasma donors, despite regulations prohibiting the use of such donors and despite
9 knowledge that the virus that causes HCV was a blood-borne disease prevalent in such
10 populations. Defendants continued using plasma taken from high-risk prison donors, even after
11 promising the FDA that they would cease doing so. Through their trade associations, Defendants
12 actively conspired to conceal these practices and to substantially delay product recalls and
13 implementation of safety measures.

14 3. Defendants failed to fully and completely disclose the known risks of their
15 products, including the risk of HCV; failed to implement readily available screening tests that
16 would have prevented HCV by excluding contaminated plasma; failed to use available methods
17 of treating plasma to kill viruses, including treatment with solvents and/or detergents; and
18 concealed and affirmatively misrepresented the extent of the health dangers of the diseases caused
19 by the products. Defendants also continued to sell old stocks of product that had not been treated
20 even after introducing a safer treated product, including stocks that Defendants knew or had
21 reason to know were made from pooled blood contaminated with HCV.

22 4. Defendants' efforts to maximize profits came at the expense of the health
23 and lives of thousands of people with hemophilia in the United States and worldwide who were
24 needlessly infected with HCV, including LOUIS FUENTES.

25 **II. JURISDICTION AND VENUE**

26 5. Plaintiff alleges an amount in controversy in excess of \$75,000, exclusive
27 of interest and costs. This Court has jurisdiction over this action pursuant to 28 U.S.C. § 1332
28 because there is complete diversity of citizenship between Plaintiff and Defendants.

1 6. Plaintiff is informed and believes and on such information and belief
2 alleges that the conduct by Defendants that is relevant to the subject matter of this action took
3 place primarily in their respective headquarters location and in other facilities within the States of
4 California and Illinois, giving these states significant contacts to the claims asserted by Plaintiff
5 and creating state interests such that the choice of either or each of these states' laws to govern
6 the adjudication of this action is neither arbitrary nor fundamentally unfair.

7 **III. PARTIES**

8 7. Plaintiff LOUIS FUENTES, a resident of Junction, Texas, who has
9 hemophilia. Plaintiff has already provided Defendant with a confidential Preliminary Patient
10 Profile Form ("PPPF"), with beginning Bates number L-PPF 003600. The PPPF contains
11 substantial additional information regarding Plaintiff's claim.

12 8. Plaintiff was infected with HCV and experienced physical and emotional
13 harm as a direct and proximate result of his use of Defendants' blood products.

14 9. Plaintiff would not have chosen to be treated with Defendants' blood
15 products, nor would have his guardians, had they known of or been informed by Defendants of
16 the true risks of using those products or the nature of the sources of the products.

17 10. Defendant CUTTER BIOLOGICAL ("CUTTER"), the predecessor of
18 Miles, Inc., and Defendant BAYER, was a California corporation headquartered in Berkeley,
19 California at all pertinent times. At all pertinent times CUTTER and its successors Miles, Inc.
20 and BAYER regularly and systematically engaged in the harvesting and collection of human
21 plasma and the processing, manufacturing, marketing, sales and distribution of factor
22 concentrates produced from such plasma, to which Plaintiff was exposed and which contributed
23 directly or indirectly to Plaintiff's infection with HCV.

24 11. Defendant BAYER CORPORATION ("BAYER"), formerly Miles, Inc., is
25 and was an Indiana corporation, authorized to do business in all 50 states and the District of
26 Columbia. Miles, Inc. had its principal place of business operation in Elkhart, Indiana, while its
27 successor BAYER has its principal place of business in Pennsylvania, with offices located at 100
28 BAYER Road, Pittsburgh, Pennsylvania 15205. At all pertinent times BAYER and its

1 predecessors Miles, Inc., and CUTTER regularly and systematically engaged in the harvesting
2 and collection of human plasma and the processing, manufacturing, marketing, sales and
3 distribution of factor concentrates produced from such plasma, to which Plaintiff was exposed
4 and which contributed directly or indirectly to Plaintiff's infection with HCV.

5 12. Defendant BAXTER HEALTHCARE CORPORATION ("BAXTER") is a
6 Delaware corporation, authorized to do business in all 50 states and the District of Columbia, with
7 its principal place of business in Illinois, with offices located at One Baxter Parkway, Deerfield,
8 Illinois 60015. At all times pertinent, Defendant BAXTER, and/or its HYLAND DIVISION, had
9 its main manufacturing plant in Glendale, California. At all times pertinent, Defendant
10 BAXTER, and/or its HYLAND DIVISION, and/or its wholly owned subsidiaries Travenol
11 Laboratories, regularly and systematically engaged in the harvesting and collection of human
12 plasma and the processing, manufacturing, marketing, sale and distribution of FACTOR
13 CONCENTRATE products produced from such plasma, to which Plaintiff was exposed and
14 which contributed directly or indirectly to Plaintiff's infection with HCV.

15 13. Defendant ARMOUR PHARMACEUTICAL COMPANY, INC.
16 ("ARMOUR") is a Delaware corporation, with its principal place of business in Pennsylvania,
17 with offices located at 500 Arcola Road, P.O. Box 1200, Collegeville, Pennsylvania, 19426-0107.
18 At all times pertinent, ARMOUR regularly and systematically engaged in the harvesting and
19 collection of human plasma and the processing, manufacturing, marketing, sales and distribution
20 of factor concentrate products produced from such plasma, to which Plaintiff was exposed and
21 which contributed directly or indirectly to Plaintiff's infection with HCV.

22
23 14. Defendant ALPHA THERAPEUTIC CORPORATION ("ALPHA") is a
24 California corporation, with its principal place of business in California, with offices at 5555
25 Valley Boulevard, Los Angeles, California 90032. At all times pertinent, Defendant has been
26 regularly and systematically engaged in the harvesting and collection of human plasma, and the
27 processing, manufacturing, marketing, sale and distribution of factor concentrate products
28

1 produced from such plasma, to which Plaintiff was exposed and which contributed directly or
2 indirectly to Plaintiff's infection with HCV.

3 15. Defendants CUTTER, BAXTER, ARMOUR and ALPHA (hereinafter
4 collectively referred to as "Defendants"), acting on behalf of themselves and/or their predecessor
5 and/or successor corporations, collected, harvested and/or processed human plasma and/or
6 manufactured, marketed, sold and distributed factor concentrate products that were contaminated
7 with HCV. In the alternative, one or more of said Defendants participated in the collection,
8 harvesting and/or processing of human plasma, and/or the manufacturing, marketing, distribution
9 and sale of factor concentrate products, that were contaminated with HCV; or assumed or became
10 responsible for, the liabilities of the Defendants and their predecessor or successor corporations
11 who did participate in the collection, harvesting and/or processing of human plasma, and/or the
12 manufacturing, marketing, distribution or sale of factor concentrate products, that were
13 contaminated with HCV, without limitation thereto.

14 16. At all times herein mentioned, Defendants were fully informed of the
15 actions of their agents and employees, and thereafter no officer, director or managing agent of
16 Defendants repudiated those actions, which failure to repudiate constituted adoption and approval
17 of said actions, and Defendants thereby ratified those actions.

18 **IV. FACTUAL ALLEGATIONS**

19 **A. Hemophilia and Its Treatment**

20 17. Hemophilia is an inherited condition that causes uncontrolled
21 hemorrhaging or bleeding. Hemophilia results from a deficiency of blood components essential
22 for coagulation. The most common form of the disease is hemophilia A, characterized by a lack
23 of a blood protein known as Factor VIII, which affects approximately one in 10,000 males.
24 Factor VIII is commonly called "AHF" or anti-hemophilic factor. Hemophilia B is characterized
25 by absence of another blood protein, known as Factor IX, affecting about one in 40,000 males.
26 Plaintiff LOUIS FUENTES has severe hemophilia A.

27 18. The treatment of hemophilia involves intravenous introduction, called
28 infusion, of the missing blood proteins required to stop bleeding. The two most prevalent forms

1 of such treatment are cryoprecipitate and factor concentrates. Factor concentrates are the
2 products made by Defendants in this action. Cryoprecipitate is made by freezing plasma, the
3 fluid component of circulating blood in which various proteins, including Factor VIII and
4 Factor IX, are contained; thawing the frozen plasma; and isolating Factor VIII from the plasma
5 through centrifugal concentration. Cryoprecipitate is an effective therapeutic agent for patients
6 with hemophilia A. Hemophilia B has been effectively treated with the use of fresh frozen
7 plasma containing Factor IX. Cryoprecipitate and fresh frozen plasma are made from small
8 numbers of donors, who are generally unpaid volunteers.

9 19. In the late 1960s to early 1970s, Defendants began to market factor
10 concentrates, which contained Factor VIII and Factor IX in higher concentrations than had been
11 available in either cryoprecipitate or fresh-frozen plasma. To produce factor concentrates,
12 Defendants mixed pools of plasma from five to over twenty thousand donors at a time, a large
13 percentage of which were paid donors. These large pools were then subjected to processes to
14 concentrate Factors VIII and IX.

15
16 **B. Defendants Failed to Disclose or Warn of Serious Adverse Effects Associated**
17 **with Factor Concentrates**

18 20. Shortly after the initial commercial marketing of Factor VIII and IX
19 concentrates in the late 1960s to early 1970s, a wide range of serious adverse effects were
20 reported in association with these products. By that time, Defendants knew of serious diseases
21 caused by unidentified agents transmissible by blood and Factor VIII and IX. Defendants failed
22 to warn Plaintiff or the medical community of these adverse effects, violating industry standards
23 and federal regulations.

24 21. By 1976, only a few years after Defendants' factor concentrate products
25 went on the market, the United States Food and Drug Administration ("FDA") Bureau of
26 Biologics held a conference titled *Unsolved Therapeutic Problems in Hemophilia*. The research
27 articles compiled from the conference discussed the high incidence of disorders in patients using
28 Defendants' products, such as liver dysfunction, enlarged spleen, Hepatitis B, and Non-A, Non-B
Hepatitis ("NANB Hepatitis," later renamed Hepatitis C). The articles concluded that these

1 disorders were tied to the patients' use of factor concentrates, and emphasized the risks entailed in
2 producing such concentrates using plasma from paid donors. For instance, Robert Gerety of the
3 FDA Bureau of Biologics, Division of Blood and Blood Products, reported that the agent or
4 agents of NANB Hepatitis "appear to be blood borne, perhaps to be associated with a form of
5 chronic hepatitis, and to represent a considerable risk to recipients who repeatedly require the
6 administration of blood products." Gerety, et al., *Viral Antigens and Antibodies in People with*
7 *Hemophilia* (1977). Gerety noted that "[t]he use of large plasma pools from paid donors no
8 doubt contributes to the risk of HBV [Hepatitis B] infection from these products," and stated that
9 "an all voluntary blood donor system is being pursued as a result of the known increased risk of
10 PTH [post-transfusion hepatitis] from blood derived from commercial donors." As described
11 below, however, Defendants not only refused to implement such a voluntary donor system, but
12 instead recruited paid donors precisely because their hepatitis exposure resulted in plasma from
13 which Defendants could make other commercially valuable products as well.

14 22. At all times material to this Complaint, Defendants failed to adequately
15 warn Plaintiff or his physicians of the serious adverse side effects of their products. Although
16 Defendants' package inserts mentioned a risk that plasma "may" contain the causative agent of
17 viral hepatitis, this warning was seriously deficient in that: (a) Defendants failed to disclose that
18 the risk of hepatitis was essentially a 100% guarantee due to their practices of using high-risk
19 donors and specifically recruiting for donors who had previously been exposed to Hepatitis B; (b)
20 while "hepatitis" simply means inflammation of the liver, and may be a relatively benign,
21 temporary condition, Defendants failed to warn that some forms of hepatitis transmitted by their
22 products were believed to present a considerable risk of severe liver damage and a significantly
23 elevated risk of liver cancer; (c) Defendants misleadingly stated that the source plasma used in
24 preparation of their products had been found to be non-reactive for Hepatitis B surface antigen
25 (HBsAg)—implying that no viral hepatitis was present in the plasma—and falsely stated that
26 available methods were not sensitive enough to detect all units of potentially infectious plasma,
27 failing to disclose that in fact Defendants had refused to implement the more sophisticated
28 Hepatitis B Core antibody (HBc) test which would have excluded the majority of plasma

1 contaminated by hepatitis; and (d) Defendants' labeling disclosed that their products were made
2 from large pools of fresh human plasma, but failed to disclose that paid donors increased the risk
3 of disease, and that the particular groups of paid donors targeted by Defendants were known to be
4 the highest risk groups.

5 23. The demand for and supply of anti-hemophilic factor rapidly increased
6 during the 1970s, with commercially-manufactured concentrate accounting for a large proportion
7 of the increase in supply. In 1977, a federal report projected that the volume of factor
8 concentrates manufactured would increase substantially by 1980. Division of Blood Diseases and
9 Resources, National Heart, Lung and Blood Institute, *Study to Evaluate the Supply-Demand*
10 *Relationships for AHF and PTC Through 1980*, at page 8; hereinafter "NHLBI Report."

11 24. In order to sell more factor concentrates to this growing market,
12 Defendants turned to the fastest and cheapest way of obtaining sufficient plasma, paid donors.
13 Defendants recruited paid donors from those populations most likely to respond to the financial
14 incentive to donate: poor inner city residents, drug abusers, prisoners, and residents of
15 impoverished developing countries such as Haiti and Nicaragua.

16 25. Defendants purposefully sought out paid donors despite knowing that the
17 risk of diseases transmissible by blood was far greater among paid donors than among volunteers.
18 Because no test was yet available in the 1970s for the NANB Hepatitis virus, an essential means
19 to prevent the virus from contaminating the plasma supply was to exclude donors with behaviors
20 that were inconsistent with good health—precisely those populations from which Defendants
21 were recruiting paid donors. Some studies indicated that paid donors were up to ten times more
22 infectious than volunteer donors. For this reason, the National Blood Policy, adopted by the
23 federal government in July 1973, advocated conversion to an all-volunteer blood supply.
24 Defendants, however, not only continued to use paid donors, but also focused their recruiting
25 efforts on the highest risk populations.

26 26. Defendants had an additional financial incentive for recruiting paid donors.
27 Factor VIII and Factor IX are only two of many products that can be made for commercial sale
28 from human plasma. According to the NHLBI Report, by the late 1970s at least 17 different

1 therapeutic components of blood were manufactured by the process of "fractionating" plasma into
2 its various elements. The NHLBI Report noted that, "as the costs of fractionation have increased,
3 fractionators have produced as many products as possible from a liter of plasma." *Id.* at 65.

4 27. Blood derivatives used as vaccines or therapeutics had particularly high
5 economic value for Defendants. The NHLBI Report noted that plasma with a very high titer, or
6 antibody level, for a corresponding antigen is "very expensive." *Id.* at 41. Such products are
7 manufactured from source plasma drawn from donors who have been sensitized to a particular
8 antigen. *Id.* The NHLBI Report specifically stated, however, that "plasma collected for high
9 antibody titer cannot be used for fractionation into therapeutic products," such as Defendants'
10 factor concentrate. *Id.* (emphasis added).

11 28. Defendants targeted donors with high titers to Hepatitis B antigens in order
12 to manufacture and sell Hepatitis B immunoglobulin (HBIG), a product that confers temporary
13 immunity to the Hepatitis B virus. Despite the warning in the NHLBI report, Defendants used the
14 same high-titer plasma obtained for making HBIG to manufacture their Factor VIII and IX
15 products used by people with hemophilia. Defendants thus sought to maximize profits by
16 producing "as many products as possible from a liter of plasma," while ignoring industry
17 standards that precluded the use of high-titer plasma for other therapeutic products.

18 29. Beginning in about 1978, Defendants began targeting homosexual donors
19 in known urban gay communities. Because urban homosexuals had been reported in the 1970s to
20 have exceptionally high prevalence of Hepatitis B infection, Defendants knew that such donors
21 would provide a reliable source of plasma for the manufacture of commercially valuable HBIG.

22 30. By the 1970s, it was also well-known in the public health community that
23 urban homosexuals engaged in promiscuous sexual practices that rapidly transmitted other
24 diseases, including NANB Hepatitis, which were transmitted by blood and were believed to have
25 serious adverse consequences. Despite this knowledge, Defendants used the same plasma pool
26 from urban homosexuals to manufacture both HBIG and Factor VIII and IX.

27 31. By the 1970s, it was also well-established that plasma from prison
28 populations carried a high risk of hepatitis and other blood-borne diseases, primarily because of

1 the concentration of intravenous (IV) drug users in prisons. By 1974, the alanine
2 aminotransferase ("ALT") test was available to test for elevated levels of liver enzymes called
3 SGOT that indicate the presence of hepatitis. Prisoners were associated with SGOT levels of
4 over 60 IUs per ml, a level that increases the risk of Hepatitis C transmission by a factor of 6.
5 Despite knowledge of this risk, Defendants actively recruited prisoners for plasma used to
6 manufacture Factor VIII and IX, while concealing or failing to disclose the risk to Plaintiff, his
7 physicians, or the FDA.

8 32. In light of Defendants' special knowledge of the disease patterns among
9 urban homosexuals and prisoners, and their recruitment of such donors for Factor VIII and IX
10 manufacture, Defendants had duties to: (a) discontinue the practice of using such high risk
11 donors; (b) disclose the risk to Plaintiff, his physicians, and the FDA, including the ongoing risk
12 of continuing to use Factor VIII and IX previously manufactured with high risk plasma and still
13 marketed to patients; (c) implement procedures to kill blood-borne diseases in the products; and
14 (d) recall existing products from distribution or further use. Instead, Defendants continued to
15 conceal their recruitment of high-risk donors and to resist warnings and recalls, and failed to
16 implement procedures to make their products safe.

17 33. By no later than 1978, Defendants knew of the availability of a new test to
18 determine whether an individual had a history of viral hepatitis, which would have disqualified
19 the donor from providing plasma for the manufacture of Factor VIII or IX. By testing a person's
20 serum for the presence of the core to the Hepatitis B antibody, a history of viral hepatitis could be
21 verified. This was known as the "HBc test." Published, peer-reviewed literature shows that the
22 HBc test was in use by researchers to determine that homosexual AIDS victims had a history of
23 viral hepatitis by no later than December 1981. Gottlieb, et al., *Pneumocystis Carinii Pneumonia*
24 *and Mucosal Candidiasis in Previously Healthy Homosexual Men*, 305 New Eng. J. Med. 1425-
25 1431 (1981).

26 34. Use of the HBc test would have eliminated approximately 75% of
27 homosexual plasma donors and over 90% of promiscuous urban homosexuals. It would have
28 eliminated almost 100% of intravenous drug users.

1 35. Use of the HBc and ALT tests together by Defendants by 1981 would have
2 eliminated the vast majority of the transmitters of HCV from the blood and plasma pools of the
3 nation, before the height of the Hepatitis C epidemic. If Defendants had implemented this test in
4 a timely manner, Plaintiff more likely than not would not have been infected with HCV as a result
5 of factor concentrate use.

6 36. As noted below, federal regulations required plasma donors to be in good
7 health, and donors with a "history of viral hepatitis" were by definition unacceptable as blood or
8 blood plasma donors. Persons with a history of viral hepatitis were excluded not only because of
9 the risk of transmitting Hepatitis B, but because such a history indicated a lifestyle or previous
10 behavior of the prospective donor that carried the risk of transmitting other viruses in addition to
11 hepatitis. A reasonable and prudent plasma fractionator would not accept a HBc positive donor
12 and expect to be in compliance with federal regulations as of 1978.

13 37. After public reports of the first hemophilia AIDS cases in July 1982,
14 government officials urged Defendants to implement the HBc test as a "surrogate" or "marker" to
15 eliminate plasma contaminated by the transmitter of AIDS and Hepatitis C. HBc testing was also
16 strongly suggested to Defendants by the CDC at a meeting of the United States Public Health
17 Service ("PHS") on January 4, 1983. Despite this urging, Defendants continued to use
18 contaminated plasma donations that would have been excluded by the HBc test and continued to
19 conceal from Plaintiff, his physicians, and the FDA the dangerous practice of targeting donors at
20 highest risk for hepatitis. At a January 6, 1983 meeting of Defendants' trade association, the
21 Pharmaceutical Manufacturer's Association, Defendants agreed not to implement the highly
22 effective HBc donor screening, and instead opted to use ineffective donor questionnaires that did
23 little to screen out donors at high-risk for Hepatitis C transmission.

24 38. As late as December 13, 1983, years after the HBc test was available, a
25 memorandum from CUTTER's responsible head, Stephen Ojala, reporting back on a meeting
26 held by Defendants, shows that Defendants conspired to propose a "task force" to further study
27 the use of HBc as an intentional, bad faith "delaying tactic for the implementation" of the test.
28

1 **C. Defendants Also Declined to Implement Available Treatment With Solvents**
2 **and/or Detergents to Kill Blood-Borne Diseases, and Continued to Dump**
3 **Contaminated Product on the Market After Safer Product Was Available**

4 39. In the late 1970s and early 1980s, it was recognized that viruses were in all
5 factor concentrate products. Treatment with solvents and/or detergents was available at that time
6 to eliminate many of these viruses, including HCV. Defendants were required to take reasonable
7 steps to eliminate contamination, but Defendants failed to utilize these available technologies to
8 eliminate the viruses in a timely manner.

9 40. Solvent and/or detergent treatment was available to Defendants by the late
10 1970s as a simple and effective method of eliminating viruses in factor concentrate products.
11 Solvents and/or detergents effectively kill viruses such as HCV by destroying the viruses' lipid
12 envelope. This method is simpler than heat treatment, and unlike heat treatment does not
13 interfere with the Factor VIII and IX proteins needed for blood clotting.

14 41. Solvents and/or detergents were well-known, commercially available
15 products as of the 1970s, and studies in which solvent and/or detergent treatment was used to
16 disrupt viruses were published in the 1970s in peer-reviewed journals. In 1980, Dr. Edward
17 Shanbrom, a former Baxter scientist, received a patent for a detergent treatment process for viral
18 inactivation of factor concentrate. Dr. Shanbrom describes the implementation of this process as
19 "as easy as washing your hands."

20 42. After receiving the patent, Dr. Shanbrom approached Defendants about
21 implementing his method, but Defendants refused to heed Dr. Shanbrom's advice. Defendants
22 refused to even commit any resources to investigate the solvent and/or detergent method.

23 43. Defendants were notified of the successful use of organic solvents to
24 destroy lipid viruses, including NANB, in factor concentrates by the New York Blood Center
25 ("NYBC") at the National Hemophilia Federation's meeting on October 27, 1983.

26 44. In 1984, Dr. Prince and Dr. Horowitz of the NYBC published the results of
27 their successful use of the solvent detergent process in well-known medical journals. They
28

1 offered to license the process to Defendants for a reasonable fee. In 1985, the NYBC obtained a
2 license from the FDA to market a solvent detergent inactivated factor concentrate.

3 45. By March, 1984, Defendants obtained licenses to sell Factor VIII treated
4 with dry heat to inactivate viruses, and Defendants had obtained such licenses for Factor IX by
5 October, 1984. The FDA did not allow them to label these products as hepatitis safe. By fall of
6 1984, Defendants were notified by treaters that previously-untreated patients in their clinical trials
7 using their dry heated products developed elevated ALT enzymes, indicative of NANB
8 infections.

9 46. Defendants were therefore aware in 1984 that dry heat did not effectively
10 inactivate the virus that causes HCV, and that solvent detergent treatment methods did eliminate
11 the risk of HCV infection, but chose not to employ the effective and efficient solvent detergent
12 technology. Instead, Defendants continued to sell their contaminated dry heat product for at least
13 four more years, resulting in the needless infection of Plaintiff and many other hemophiliacs.

14 47. A recent CDC study documented the comparative effectiveness of the dry
15 heat and solvent detergent inactivation methods. The study reported that "84% of previously
16 untreated patients infused with dry-heated Factor VIII products developed non-A, non B
17 hepatitis..." Soucie, Richardson, Evatt et al., *Risk Factor for Infection with HBV and HCV in a*
18 *Large Cohort of Hemophiliac Males*, 41 Transfusion 338-343 (2001).

19 48. The same CDC study reported that "solvent detergent treatment of blood
20 components [was] found to be more effective against enveloped viruses than heat treatment ...
21 No cases of HBV, HCV, or HIV transmission through solvent detergent virus inactivated
22 products have been found in prospective studies of previously untreated patients..."

23 49. The study further reported "in our data, the first dramatic decline in HCV
24 prevalence appears in the 1987 birth cohort. The drop in HCV transmission correlates with the
25 licensing of solvent detergent treatment of Factor IX products in 1987. In addition, this cohort
26 would have been the first to benefit from the screening of blood donors using the surrogate
27 markers ALT (begun in late 1986) and anti-HBc (begun in 1987), testing that was associated with
28 a markedly decreased risk of HCV infection from blood transfusions."

1 50. The study states further that “the residual transmissions after 1987 possibly
2 represent the use of product already manufactured or product manufactured during the interval
3 required to implement the new technology. The 18-month shelf life of factor concentrates placed
4 those people with hemophilia born as late as 1989 at risk of infection.” The study goes on to
5 recommend testing for all people with hemophilia who received infusions of Defendants’ blood
6 products prior to 1992.

7 51. By 1988, it was clear to the medical and scientific community what
8 Defendants had long known: dry-heated factor concentrates were transmitting the potentially
9 deadly NANB virus, and safer products were available. This knowledge prompted the CDC to
10 publish recommendations that dry-heated products no longer be used by hemophiliacs.
11 Defendants continued sales of their dry-heated products after these warnings, however, and never
12 undertook a large-scale recall of dry-heated product. Defendants finally introduced solvent
13 detergent-treated products to the market in 1988 and 1989, but continued to sell their NANB-
14 contaminated dry-heated factor concentrates after this date.

15 52. The failure of Defendants to implement solvent and/or detergent viral
16 inactivation techniques in a timely manner, to warn of the risk that dry heat treated Factor VIII
17 and IX blood products could transmit HCV, and to recall dry heat-treated products that posed this
18 risk caused the needless infection of thousands of people with hemophilia with HCV, including
19 Plaintiff. Even after Defendants knew or should have known that solvent and/or detergents
20 effectively destroyed HCV, they continued to sell dry heat-treated Factor VIII and IX, and
21 refused to recall these dangerous products from the market.

22
23 D. **Defendants Fraudulently Misrepresented the Safety, and Concealed the**
24 **Dangers, of Their Factor VIII and IX Products**

25 53. Defendants engaged in a pattern and practice of fraudulent concealment of
26 their dangerous practices, fraudulent misrepresentations regarding their efforts to assure safety,
27 and fraudulent misrepresentations regarding the risk of Hepatitis C, in order to maintain profits
28 from both factor concentrates and HBIG. A summary of Defendants’ fraudulent
misrepresentations and concealment is set forth below.

1 54. On July 27, 1982, a meeting of the Public Health Service was held as the
2 result of the CDC's report that three people with hemophilia had contracted AIDS. The
3 responsible heads of Defendants were in attendance, along with officials from the National
4 Hemophilia Foundation, CDC and FDA. Defendants were aware that they had used plasma from
5 known, targeted homosexuals in the manufacture of their Factor VIII and IX blood products.
6 These products had a shelf life of two years and were either in production or already on the
7 shelves in pharmacies waiting to be infused by people with hemophilia who purchased them.
8 Defendants failed to disclose these facts at the meeting where CDC officials were present, despite
9 knowledge that the CDC's primary concern at that meeting was the contamination of Factor VIII
10 and IX by the agent that transmitted AIDS, which, like hepatitis, was already well-known to be
11 epidemic in the targeted homosexual population. (CUTTER memorandum dated August 3,
12 1982.)

13 55. In or about December, 1982, Rodell, the responsible head for BAXTER,
14 entered into an agreement with officials of the FDA to the effect that BAXTER would no longer
15 use prison plasma in the production of factor concentrates. In fact, BAXTER, unbeknownst to
16 the FDA, continued to use prison plasma in factor concentrate production through October 1983.
17 BAXTER memorandum dated October 20, 1983.

18 56. On January 5, 1983, an AIDS meeting was held at Children's Orthopedic
19 Hospital in Los Angeles, California, the largest hemophilia treatment center in the United States.
20 Representatives of Defendants were present at the meeting with treaters and patients. A patient
21 asked representatives from Defendants the following question: "Is the plasma from homosexuals,
22 prisoners, Haitians or other high risk persons being used in the manufacture of concentrates?"
23 Defendants did not admit targeting or using plasma from homosexuals, prisoners or inner city IV
24 drug abusers. Defendants' representatives made no response to the question, thereby concealing
25 the true risk created by the use of plasma from known homosexuals, IV drug abusers and
26 prisoners in the manufacture of factor concentrates.

27 57. At the January 5, 1983 meeting, and in the presence of the patients, one of
28 the treating physicians, Dr. Kasper, asked CUTTER's Stephen Ojala: "These [plasma] centers

1 seem to be in rundown centers of town. Is there a move to move them to rural towns?" Ojala
2 answered: "Many of the centers are in smaller communities and in towns such as Ypsilanti,
3 Seattle, Clayton, NC., and San Diego. We do not have centers in L.A. or San Francisco." This
4 answer was misleading because Ojala failed to state that CUTTER's largest and first plasma
5 center was located at Arizona State Penitentiary. CUTTER also had a center at the Las Vegas
6 Prison. Ojala and CUTTER were well aware of the CDC's and FDA's concern over use of prison
7 plasma, due to homosexual practices and drug abuse in the prison donor population. Many of
8 CUTTER'S centers were in inner city areas frequented by IV drug abusers, such as downtown
9 Oakland, California. CUTTER had also used plasma from centers which targeted known
10 homosexuals. In August 1982, CUTTER quarantined plasma from the Valley Medical Center, a
11 center which targeted known homosexuals, because a donor was hospitalized with full blown
12 AIDS. The plasma was intended for factor concentrate and HBIG production, but was not used
13 because it had thawed on the way to the processing plant. Upon receiving a report of this incident
14 from CUTTER, the FDA indicated a recall might have been necessary if the plasma had been
15 incorporated into factor concentrate final product. Ojala omitted any mention of these facts and
16 circumstances in his response to Dr. Kasper regarding the location of their plasma centers.
17 (CUTTER memorandum dated January 5, 1983.)

18 58. On January 14, 1983, responsible heads from Defendants attended a
19 meeting of the National Hemophilia Foundation ("NHF"). Defendants were very concerned that
20 the NHF would insist on a recommendation that HBc testing be implemented, consistent with the
21 CDC recommendation 10 days earlier. In order to defer a NHF recommendation that HBc testing
22 be used, Michael Rodell, a representative of BAXTER, told NHF officials on behalf of
23 Defendants, that surrogate testing was in the "R and D," or "Research and Development," stage
24 currently. Rodell concealed the fact that the CDC had strongly recommended use of the HBc
25 antibody test as a screening device for high risk donors. The HBc antibody test was not in the "R
26 and D" stage, and was suitable for use as a screening device for high risk AIDS and Hepatitis C
27 donors. In fact, the HBc test had been approved in 1979 by the FDA as a test to be used to
28 ascertain a history of previous hepatitis B infection, and to screen blood and plasma donors.

1 Donors with a hepatitis history were specifically prohibited pursuant to the federal regulations (21
2 C.F.R. § 640.63). Rodell acknowledged that implementation of the HBc test would eliminate
3 high titered immunoglobulin donors, but failed to disclose that opposition to use of the test was
4 based on economic rather than safety concerns.

5 59. At the January 14, 1983 meeting, Defendants concealed their advertising in
6 publications distributed among urban homosexuals, for the specific purpose of attracting them to
7 plasma centers which supplied high titered plasma to Defendants. Defendants also concealed
8 their extensive use of prison plasma, and failed to reveal their "gentlemen's agreement" with the
9 FDA to discontinue use of these plasma sources immediately. (CUTTER Memorandum dated
10 January 17, 1983.)

11 60. On or about December 15, 1983, Rodell, then the head of Armour
12 Pharmaceutical Company, Inc., told members of the federal Blood Product Advisory Committee
13 (BPAC) and FDA officials that Defendants wanted a three-month deferral in implementation of
14 any recommendations by the BPAC or FDA that HBc testing be required for plasma donors.
15 Rodell told the FDA that the purpose of the deferral was to prepare a response to the proposed
16 recommendation. In fact, Defendants had agreed to seek the three-month hiatus as a "delaying
17 tactic" to avoid implementing the test, and the request for a deferral was made in bad faith.
18 (CUTTER memorandum dated December 13, 1983.)

19 61. Defendants fraudulently misrepresented the risk of Hepatitis C due to
20 factor concentrates, failed to disclose accurate warnings of the risk to Plaintiff or his physicians,
21 and fraudulently purported to be doing "everything possible" to improve safety, when in fact
22 Defendants maximized the risk by recruiting high-risk donors and by resisting and obstructing
23 HBc testing, treatment with solvents and/or detergents, and other measures that would truly have
24 reduced the risk.

25 **E. Defendants' Activities Were Subject to Applicable Federal Regulations,**
26 **Which Evidence the Standard of Care With Which Defendants Should Have**
27 **Complied**

28 62. Blood derivatives such as Factor VIII and IX are prescription biologicals
subject to federal regulation as both "biological products" and "drugs." Public Health Service

1 Act, "Regulation of Biological Products," 42 U.S.C. § 262; Food, Drug & Cosmetic Act
2 ("FDCA"), 21 U.S.C. § 301, *et seq.* (2005).

3 (a) 21 U.S.C. § 331(b) prohibited and continues to prohibit
4 "adulteration or misbranding of any ... drug"

5 (b) 21 U.S.C. § 351(a)(2)(B) provided and continues to provide that
6 "[a] drug ... shall be deemed to be adulterated ... if ... the methods used in, or the facilities or
7 controls used for, its manufacture, processing, packing, or holding do not conform to or are not
8 operated or administered in conformity with current good manufacturing practice to assure that
9 such drug meets the requirements of this chapter as to safety. ..."

10 (c) 21 U.S.C. § 352 provided and continues to provide that "[a] drug...
11 shall be deemed to be misbranded. ... if its labeling is false or misleading in any particular."

12 (d) 21 U.S.C. § 352(f)(2) provided and continues to provide that a drug
13 shall be deemed to be "misbranded" unless its labeling bears "adequate warnings against use. ...
14 where its use may be dangerous to health."

15 (e) 21 U.S.C. § 352(n) provided and continues to provide that a drug
16 shall be deemed to be "misbranded" unless the labeling included information concerning side
17 effects and contraindications as required in federal regulations.

18 (f) 21 U.S.C. § 321(n) provided and continues to provide that if an
19 article is alleged to be misbranded because the labeling or advertising is misleading, then the
20 determination of whether the labeling or advertising is misleading shall take into account "not
21 only representations made or suggested" by affirmative statements, "but also the extent to which
22 the labeling or advertising fails to reveal facts material in the light of such representations or
23 material with respect to consequences which may result from the use" of the drug.

24 63. At all times material to this Complaint, 21 C.F.R. § 201.57(e) provided and
25 continues to provide as follows, with respect to information to be provided with the sale of
26 Defendants' products:

27 Warnings: Under this section heading, the labeling shall describe
28 serious adverse reactions and potential safety hazards, limitations in
use imposed by them, and steps that should be taken if they occur.
The labeling shall be revised to include a warning as soon as there

1 is reasonable evidence of an association with a drug; a causal
2 relationship need not have been proved.

3 64. At all times material to this Complaint, 21 C.F.R. § 200.5 provided and
4 continues to provide as follows:

5 Manufacturers and distributors of drugs and the Food and Drug
6 Administration occasionally are required to mail important
7 information about drugs to physicians and others responsible for
8 patient care. In the public interest, such mail shall be distinctive in
9 appearance so that it will be promptly recognized and read.

10 65. At all times material to this Complaint, Part 606 of 21 C.F.R. set forth and
11 continues to set forth "Current Good Manufacturing Practices" for biological products generally,
12 and 21 C.F.R. § 640, *et seq.*, set forth additional good manufacturing practices for blood and
13 plasma biologicals.

14 66. At all times material to this Complaint, 21 C.F.R. § 606.140(a) provided
15 and continues to provide:

16 Laboratory control procedures shall include: The establishment of
17 scientifically sound and appropriate specifications, standards and
18 test procedures to assure that blood and blood components are safe,
19 pure, potent and effective.

20 67. At all times material to this Complaint, 21 C.F.R. § 640.60 defined and
21 continues to define "Source Plasma" as:
22 the fluid portion of human blood collected by plasmapheresis, and
23 is intended as source material for further manufacturing use.

24 68. At all times material to this Complaint, 21 C.F.R. § 640.63(c), (1999),
25 titled "Qualification of Donor," provided and continues to provide as follows with respect to
26 donors of source plasma:

27 Donors shall be in good health on the day of donation, as indicated
28 in part by: . . . (9) freedom from any disease, other than malaria,
transmissible by blood transfusion, in so far as can be determined
by history and examination indicated in this section; (10) freedom
of the arms and forearms from skin punctures or scars indicative of
addiction to self-injected narcotics; (11) freedom from a history of
viral hepatitis; (12) freedom from a history of close contact within
six months of donation with an individual having viral
hepatitis; . . .

1 Further, 21 C.F.R. § 640.63(a) provided and continues to provide that the method of determining
2 "suitability of a donor" included "tests" as well as the taking of a history and physical
3 examination.

4 69. The foregoing statutes and regulations are evidence of the standard of care
5 Defendants should have employed in the manufacture and sale of Factor VIII and Factor IX.
6 Defendants violated the foregoing regulations and/or failed to comply with applicable standards
7 of care by: (a) marketing "adulterated" products that were unsafe as a result of failure to comply
8 with "Current Good Manufacturing Practice"; (b) marketing "misbranded" products that were
9 misleading and failed to disclose or warn of health dangers; (c) failing to warn of serious adverse
10 reactions and potential safety hazards as soon as there was reasonable evidence of an association
11 with their products; (d) failing to exclude intravenous drug users who were unsuitable donors;
12 (e) failing to exclude donors with a history of viral hepatitis who were unsuitable donors;
13 (f) affirmatively seeking out unsuitable donors known to have viral hepatitis antibodies, as well as
14 prison populations known to include substantial numbers of intravenous drug users, for inclusion
15 of their plasma in the pools used to make Factor VIII and Factor IX; (g) failing to disclose their
16 use of dangerous donors; and (h) failing to use appropriate tests and/or procedures to assure their
17 products were safe.

18 **F. Group Liability**

19 70. All Defendants likely to have caused the harm to Plaintiff are parties to this
20 lawsuit and properly before the court.

21 71. The conduct of Defendants, with respect to their Factor VIII and Factor IX
22 products and related plasma collection methods, was tortious.

23 72. The harm which has been caused to Plaintiff resulted from the conduct of
24 one, or various combinations of the Defendants, and, through no fault of Plaintiff, there may be
25 uncertainty as to which one or combination of Defendants caused the harm.

26 73. The burden of proof should be upon each Defendant to prove that the
27 Defendant has not caused the harms suffered by the Plaintiff.

28 74. Factor concentrates were manufactured using the same fractionation

1 method by all Defendants. As such, during the relevant years, factor concentrates were a fungible
2 product, and physicians prescribed the products interchangeably without regards to brand names.

3 75. The factor concentrates manufactured by Defendants contained the same
4 design flaws. They were all manufactured from paid donor plasma, which was at highest risk for
5 Hepatitis B and Hepatitis C viral transmission. In addition, all Defendants' factor concentrates
6 were made from large pools consisting of 5,000 to over 20,000 paid donors, which further
7 magnified the risk of viral transmission.

8 76. None of the factor concentrates made by Defendants during the relevant
9 time period were subjected to viral inactivation processes such as solvent and/or detergent
10 treatment that were effective against HCV. Therefore, all of Defendants' factor concentrates
11 carried a significant risk of HCV transmission during this time. In addition, all of Defendants'
12 factor concentrate products were similarly misbranded. All of the products failed to warn of the
13 known risks enumerated in this complaint.

14 **V. TOLLING OF APPLICABLE STATUTES OF LIMITATION**

15 77. Any and all potentially applicable statutes of limitations have been tolled
16 by Defendants' affirmative and intentional acts of fraudulent conduct, concealment, and
17 misrepresentation, alleged above, which estop Defendants from asserting statutes of limitation.
18 Such acts include but are not limited to intentionally covering up and refusing to disclose use of
19 high-risk plasma; selling products known to be contaminated; suppressing and subverting medical
20 and scientific research; and failing to disclose and suppressing information concerning the risk of
21 HCV transmission from Defendants' contaminated factor concentrates.

22 78. Defendants are estopped from relying on any statutes of limitation because
23 of their fraudulent concealment and misrepresentation alleged above. Defendants were under a
24 duty to disclose the precise risks of HCV transmission from their contaminated factor concentrate
25 because this is nonpublic information over which they had exclusive control, because Defendants
26 knew this information was not readily available to people with hemophilia like Plaintiff, and
27 because this information was relevant to such people in deciding whether to use Defendants'
28 factor concentrate.

79. Until very recently, Plaintiff had no knowledge that Defendants were engaged in much of the wrongdoing alleged herein. Because of the fraudulent and active concealment of the wrongdoing by Defendants, including but not limited to deliberate efforts—which continue to this day—to give Plaintiff the materially false impression that Defendants undertook all feasible safety precautions to reduce the risk of HCV transmission from their contaminated factor concentrates, Plaintiff could not reasonably have discovered the wrongdoing any time prior to this time, nor could Plaintiff have, as a practical matter, taken legally effective action given the unavailability, until very recently, of internal memoranda and other documents (as generally described herein) as evidence in support of Plaintiff's claims. Defendants still refuse to admit and continue to conceal their wrongdoing, and therefore Defendants' acts of fraudulent concealment and misrepresentation continue through the present time.

VI. CLAIMS FOR RELIEF

FIRST CLAIM FOR RELIEF

FRAUDULENT OMISSION AND CONCEALMENT

80. Plaintiff incorporates by reference all previous paragraphs of this Complaint as if fully set forth here and further alleges as follows:

81. Defendants had a confidential and special relationship with Plaintiff due to: (a) Defendants' vastly superior knowledge of the health and safety risks relating to Factor VIII and Factor IX; (b) Defendants' sole and/or superior knowledge of their dangerous and irresponsible plasma collection practices; and (c) Defendants' direct communications with the hemophiliac community through newsletters that purported to accurately convey the risk of NANB. As a result, Defendants had an affirmative duty to fully and adequately warn the hemophiliac community, including Plaintiff, his guardians, and his physicians, of the true health and safety risks related to their Factor VIII and Factor IX blood products and constituent plasma, and a duty to disclose their dangerous and irresponsible plasma collection practices. Independent of any special relationship of confidence or trust, Defendants had a duty not to conceal the dangers of their products to Plaintiff, his guardians, and his physicians.

1 86. As a direct and proximate result of Defendants' fraudulent concealment
2 and suppression of material health and safety risks relating to their Factor VIII and Factor IX
3 blood products and of Defendants' dangerous and irresponsible plasma collection practices,
4 Plaintiff has suffered and will continue to suffer injury, harm and economic loss. As the direct,
5 proximate and legal result of the Defendants' fraudulent concealment and suppression of material
6 health and safety risks relating to their Factor VIII and Factor IX blood products and of
7 Defendants' dangerous and irresponsible plasma collection practices, Plaintiff has been injured
8 and has incurred damages, including but not limited to physical injuries to his person, medical
9 expenses in the past, past disability, past loss of use of the body, and past physical and mental
10 pain and suffering; and may incur in the future medical and hospital expenses, permanent
11 disability, loss of use of the body, physical and mental pain and suffering, and loss of the
12 enjoyment of life.

13 87. Plaintiff is therefore entitled to damages in an amount to be proven at trial,
14 together with interest thereon and costs.

15 88. Defendants' conduct, as alleged above, was malicious, intentional and
16 outrageous and constituted willful and wanton disregard for the rights or safety of others. Such
17 conduct was directed specifically at Plaintiff and warrants an award of punitive damages.

18 89. Plaintiff is informed and believes that Defendants utilize retention policies
19 that provide for scheduled destruction of documents and other items, which may result in the
20 knowing, negligent, or inadvertent destruction of documents, data, and materials relevant and
21 necessary to adjudication of this action, including, but not limited to, records identifying batch or
22 lot numbers of Defendants' products shipped to particular treatment facilities, which may
23 facilitate product tracing. This risk warrants an order from this Court that such evidence
24 (including all documents, data compilations, and tangible things within the meaning of Rule 26 of
25 the Federal Rules of Civil Procedure) be preserved and maintained for use in these proceedings.
26
27
28

1 82. Misrepresentations made by Defendants about the health and safety of their
 2 factor concentrate products independently imposed a duty upon Defendants to fully and
 3 accurately disclose to the hemophiliac community, including Plaintiff, his guardians, and his
 4 physicians, the true health and safety risks related to Factor VIII and Factor IX and its constituent
 5 plasma, and a duty to disclose their dangerous and irresponsible plasma collection practices.

6 83. In connection with their Factor VIII and Factor IX products, Defendants
 7 fraudulently and intentionally concealed important and material health and safety product risk
 8 information from Plaintiff, his guardians, the hemophiliac community, and treating physicians, all
 9 as alleged in this Complaint.

10 84. Any of the following is sufficient to independently establish Defendants'
 11 liability for fraudulent omission and/or concealment:

- 12 a. Defendants fraudulently concealed the health and safety hazards,
 13 symptoms, diseases and/or health problems associated with their
 14 Factor VIII and Factor IX blood products and related plasma collection
 15 activities;
- 16 b. Defendants fraudulently concealed the practice of using unsuitable plasma
 17 from unsuitable donors in the manufacture of their Factor VIII and
 18 Factor IX blood products;
- 19 c. Defendants fraudulently concealed their practice of avoiding the use of
 20 available technology to detect viruses in their Factor VIII and Factor IX
 21 blood products and the components thereof;
- 22 d. Defendants fraudulently concealed their practice of avoiding the use of
 23 available technology to destroy viruses in their Factor VIII and Factor IX
 24 blood products and the components thereof; and/or
- 25 e. Defendants fraudulently concealed information about the known
 26 comparative risks and benefits of the use of their Factor VIII and Factor IX
 27 and the relative benefits and availability of alternate products and therapies.

28 85. Defendants knew that Plaintiff, his guardians, the hemophiliac community,
 and physicians would regard the matters Defendants concealed to be important in determining a
 course of treatment, including the decision whether to use their Factor VIII and/or Factor IX
 blood products.

SECOND CLAIM FOR RELIEF

BREACH OF IMPLIED WARRANTY

90. Plaintiff incorporates by reference all previous paragraphs of this Complaint as if fully set forth here and further alleges as follows:

91. Defendants' factor concentrate products were intentionally designed, manufactured, promoted, distributed and sold to be introduced into the human body.

92. Defendants breached the implied warranties of merchantability and fitness because Defendants' factor concentrate products cannot pass without objection in the trade, are unsafe, are not merchantable, are unfit for their ordinary use when sold, and are not adequately packaged and labeled.

93. Plaintiff is therefore entitled to damages in an amount to be proven at trial, together with interest thereon and costs.

THIRD CLAIM FOR RELIEF

NEGLIGENCE

94. Plaintiff incorporates by reference all previous paragraphs of this Complaint as if fully set forth here and further alleges as follows:

95. Defendants marketed their Factor VIII and/or Factor IX blood products to and for the benefit of Plaintiff, and knew or should have known that Plaintiff would use their Factor VIII and/or Factor IX blood products.

96. Defendants owed Plaintiff duties to exercise reasonable or ordinary care under the circumstances in light of the generally recognized and prevailing best scientific knowledge.

97. Through the conduct described in the foregoing and subsequent paragraphs of this Complaint, Defendants breached their duties to Plaintiff. The following sub-paragraphs summarize Defendants' breaches of duties to Plaintiff and describe categories of acts or omissions constituting breaches of duties by Defendants. Each and/or any of these acts or omissions establishes an independent basis for Defendants' liability in negligence:

- a. Failure to exercise reasonable care in producing Factor VIII and Factor IX blood products that were free of viruses, including the virus that causes Hepatitis C;
- b. Failure to exercise reasonable care in assuring that only suitable plasma would be used in manufacturing their Factor VIII and Factor IX blood products;
- c. Failure to exercise reasonable care in testing plasma used in manufacturing their Factor VIII and Factor IX blood products for viral contamination;
- d. Failure to exercise reasonable care in recruiting and screening donors of plasma used in their manufacture of Factor VIII and Factor IX blood products;
- e. Failure to reasonably employ anti-viral techniques, including solvent and/or detergent treatment, in the manufacture of their Factor VIII and Factor IX blood products;
- f. Unreasonable overpromotion of their Factor VIII and Factor IX blood products;
- g. Understating the relative value of hemophilia treatments that constituted alternatives to their Factor VIII and Factor IX blood products;
- h. Failure to warn physicians, Plaintiff, his guardians, and the hemophilia community of the dangers associated with their Factor VIII and Factor IX blood products and/or the viruses and foreign bodies contained within the plasma used in manufacturing their Factor VIII and Factor IX blood products;
- i. Failure to exercise reasonable care by complying with federal regulations then applicable to plasma collection and the manufacture of Factor VIII and Factor IX blood products;
- j. Failure to exercise reasonable care in disseminating information about their methods of manufacturing their Factor VIII and Factor IX blood products and the risks that were created by said methods; and
- k. Failure to exercise reasonable care in recalling their Factor VIII and Factor IX blood products.

98. Defendants knew, or should have known, that due to their failure to use reasonable care, Plaintiff and other people with hemophilia would use and did use Defendants' Factor VIII and/or Factor IX products to the detriment of their health, safety and well-being.

99. As the direct, proximate and legal result of the Defendants' negligence, Plaintiff has been injured and has incurred damages, including but not limited to physical injuries

1 to his person, medical expenses in the past, past disability, past loss of use of the body, and past
2 physical and mental pain and suffering; and may incur in the future medical and hospital
3 expenses, permanent disability, loss of use of the body, physical and mental pain and suffering,
4 and loss of the enjoyment of life.

5 100. Plaintiff is therefore entitled to damages in an amount to be proven at trial,
6 together with interest thereon and costs.

7 101. Defendants' conduct, as alleged above, was malicious, intentional and
8 outrageous, and constituted willful and wanton disregard for the rights or safety of others. Such
9 conduct was directed specifically at Plaintiff and warrants an award of punitive damages.

10 **FOURTH CLAIM FOR RELIEF**

11 **NEGLIGENCE PER SE**

12 102. Plaintiff incorporates by reference all previous paragraphs of this
13 Complaint as if fully set forth here and further alleges as follows:

14 103. Defendants violated applicable federal statutes and regulations relating to
15 prescription drugs. Plaintiff is a person whom these statutes and regulations were meant to
16 protect.

17 104. Defendants' violation of these statutes or regulations constitutes negligence
18 per se.

19 105. Defendants' violation of these statutes or regulations was the direct,
20 proximate and legal cause of Plaintiff's injuries and damages. As the direct and legal result of the
21 Defendants' negligence, Plaintiff has been injured and has incurred damages, including but not
22 limited to physical injuries to his person, medical expenses in the past, past disability, past loss of
23 use of the body, and past physical and mental pain and suffering; and may incur in the future
24 medical and hospital expenses, permanent disability, loss of use of the body, physical and mental
25 pain and suffering, and loss of the enjoyment of life.

26 106. Plaintiff is therefore entitled to damages in an amount to be proven at trial,
27 together with interest thereon and costs.

107. Defendants' conduct, as alleged above, was malicious, intentional and outrageous and constituted willful and wanton disregard for the rights or safety of others. Such conduct was directed specifically at Plaintiff and warrants an award of punitive damages.

VII. PRAYER FOR RELIEF

WHEREFORE, Plaintiff prays for judgment against Defendants as follows:

108. For compensatory damages sustained by Plaintiff against Defendants in an amount to be determined at trial;

109. For punitive and exemplary damages according to proof against Defendants;

110. For an award of prejudgment interest, costs, disbursements and reasonable attorneys' fees;

111. For injunctive relief in the form of an order requiring Defendants to preserve all relevant documents; and

112. For such other and further relief as the Court deems equitable or appropriate under the circumstances.

Dated: February 22, 2008



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DEMAND FOR JURY TRIAL

Plaintiff demands a trial by jury on all issues stated.

Dated: February 22, 2008



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